- (2) The PRRB has the authority to review the action taken by HCFA on the facility's requests. However, the PRRB's decision is subject to review by the Administrator under §405.1875 of this chapter.
- (3) A facility must request and obtain a final agency decision, in accordance with paragraph (b)(1) of this section, prior to seeking judicial review of the denial, in whole or in part, of the exception request.
- (c) *Procedure.* (1) The facility must request review within 180 days of the date of the decision on which review is sought.
- (2) The facility may not submit to the reviewing entity, whether it is the intermediary or the PRRB, any additional information or cost data that had not been submitted to HCFA at the time HCFA evaluated the exception request.
- (d) Determining amount in controversy. For purposes of determining PRRB jurisdiction under subpart R of part 405 of this chapter for the appeals described in paragraph (b) of this section—
- (1) The amount in controversy per treatment is determined by subtracting the amount of program payment from the amount the facility requested under §413.180; and
- (2) The total amount in controversy is calculated by multiplying the amount in controversy per treatment by the projected number of treatments for the exception request period.

§413.196 Notification of changes in rate-setting methodologies and payment rates.

- (a) HCFA or the facility's intermediary notifies each facility of changes in its payment rate. This notice includes changes in individual facility payment rates resulting from corrections or revisions of particular geographic labor cost adjustment factors.
- (b) Changes in payment rates resulting from incorporation of updated cost data or general revisions of geographic labor cost adjustment factors are announced by notice published in the FEDERAL REGISTER without opportunity for prior comment. Revisions of the rate-setting methodology are pub-

lished in the FEDERAL REGISTER in accordance with the Department's established rulemaking procedures.

§413.198 Recordkeeping and cost reporting requirements for outpatient maintenance dialysis.

- (a) Purpose and Scope. This section implements section 1881(b)(2)(B)(i) of the Act by specifying recordkeeping and cost reporting requirements for ESRD facilities approved under subpart U of part 405 of this chapter. The records and reports will enable HCFA to determine the costs incurred in furnishing outpatient maintenance dialysis as defined in §413.170(a).
- (b) Recordkeeping and reporting requirements. (1) Each facility must keep adequate records and submit the appropriate HCFA-approved cost report in accordance with §§ 413.20 and 413.24, which provide rules on financial data and reports, and adequate cost data and cost finding, respectively.
- (2) The cost reimbursement principles set forth in this part (beginning with §413.134, Depreciation, and excluding the principles listed in paragraph (b)(4) of this section), apply in the determination and reporting of the allowable cost incurred in furnishing outpatient maintenance dialysis treatments to patients dialyzing in the facility, or incurred by the facility in furnishing home dialysis service, supplies, and equipment.
- (3) Allowable cost is the reasonable cost related to dialysis treatments. Reasonable cost includes all necessary and proper expenses incurred by the facility in furnishing the dialysis treatments, such as administrative costs, maintenance costs, and premium payments for employee health and pension plans. It includes both direct and indirect costs and normal standby costs. Reasonable cost does not include costs that—
- (i) Are not related to patient care for outpatient maintenance dialysis;
- (ii) Are for services or items specifically not reimbursable under the program;
- (iii) Flow from the provision of luxury items or servicess (items or services substantially in excess of or more

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expensive than those generally considered necessary for the provision of needed health services); or

- (iv) Are found to be substantially out of line with other institutions in the same area that are similar in size, scope of services, utilization, and other relevant factors.
- (4) The following principles of this part do not apply in determining adjustments to allowable costs as reported by ESRD facilities:
- (i) Section 413.157, Return on equity capital of proprietary providers;
- (ii) Section 413.200, Reimbursement of OPAs and histocompatibility laboratories:
- (iii) Section 413.9, Cost related to patient care (except for the principles stated in paragraph (b)(3) of this section); and
- (iv) Sections 413.64, Payments to providers, and §§ 413.13, 413.30, 413.35, 413.40, 413.74, and §§ 415.55 through 415.70, § 415.162, and § 415.164 of this chapter, Principles of reimbursement for services by hospital-based physicians.

§413.200 Payment of independent organ procurement organizations and histocompatibility laboratories.

- (a) Principle. Covered services furnished after September 30, 1978 by organ procurement organizations (OPOs) and histocompatibility laboratories in connection with kidney acquisition and transplantation will be reimbursed under the principles for determining reasonable cost contained in this part. Services furnished by freestanding OPOs and histocompatibility laboratories, that have an agreement with the Secretary in accordance with paragraph (c) of this section, will be reimbursed by making an interim payment to the transplant hospitals using these services and by making a retroactive adjustment, directly with the OPO or laboratory, based upon a cost report filed by the OPO or laboratory. (The reasonable costs of services furnished by hospital based OPOs or laboratories will be reimbursed in accordance with the principles contained in §§ 413.60 and 413.64.)
- (b) *Definitions*. For purposes of this section:

Freestanding refers to an OPO or a histocompatibility laboratory that is not—

- (1) Subject to the control of the hospital with respect to the hiring, firing, training, and paying of employees; and
- (2) Considered as a department of the hospital for insurance purposes (including malpractice insurance, general liability insurance, worker's compensation insurance, and employee retirement insurance).

Histocompatibility laboratory means a laboratory meeting the standards and providing the services for kidneys or other organs set forth in §413.2171(d) of this chapter.

OPO means an organization defined in §486.302 of this chapter.

- (c) Agreements with independent OPOs and laboratories. (1) Any freestanding OPO or histocompatibility laboratory that wishes to have the cost of its pretransplant services reimbursed under the Medicare program must file an agreement with HCFA under which the OPO or laboratory agrees—
- (i) To file a cost report in accordance with §413.24(f) within three months after the end of each fiscal year;
- (ii) To permit HCFA to designate an intermediary to determine the interim reimbursement rate payable to the transplant hospitals for services provided by the OPO or laboratory and to make a determination of reasonable cost based upon the cost report filed by the OPO or laboratory;
- (iii) To provide such budget or cost projection information as may be required to establish an initial interim reimbursement rate;
- (iv) To pay to HCFA amounts that have been paid by HCFA to transplant hospitals and that are determined to be in excess of the reasonable cost of the services provided by the OPO or laboratory; and
- (v) Not to charge any individual for items or services for which that individual is entitled to have payment made under section 1861 of the Act.
- (2) The initial cost report due from an OPO or laboratory is for its first fiscal year during any portion of which it had an agreement with the Secretary under paragraphs (c) (1) and (2) of this section. The initial cost report covers